

WHAT IS CLAIMED IS:

- A method of inhibiting inflammatory leukocyte mediated destruction of tissue in a patient, the method comprising administering to the patient a composition comprising a \$1-integrin inhibitor.
- The method of claim 1 wherein the β1-integrin inhibitor is a peptide 2. comprising a C-terminal LipAr motif.
- 3. The method of claim 2 wherein the β1-integrin inhibitor is a peptide comprising an amino acid sequence selected from the group consisting of WQPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAAIY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WQPPDADIY (SEQ ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEO ID NO:9), PRQAWRPIY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARIY (SEQ ID NO:12), RARIY (SEQ ID NO:13), ARIY (SEQ ID NO:14), and RIY.
- 4. The method of claim 1 wherein the composition further includes a pharmaceutically acceptable carrier.
- 5. The method of claim 1 wherein the inflammatory leukocyte mediated destruction of tissue occurs as a result of CNS ischemic injury, myocardial infarction, angioplasty, surgical incisions, injury-related trauma, transplant reperfusion, or a combination thereof.
- 6. The method of claim 1 wherein the inflammatory leukocyte mediated destruction of tissue occurs as a result of exposure to heat, cold, light, electricity, chemicals, or a combination thereof.

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- A method of treating a stroke patient, the method comprising administering 7. to the patient a composition comprising a β1-integrin inhibitor in an amount effective to reduce infarct size, reduce neurological deficit, or both.
- The method of claim 7 wherein the composition is administered locally. 8.
- The method of claim 8 wherein the \beta1-integrin inhibitor is administered in 9. an amount effective to reduce the infarct size by at least about 80%.
- The method of claim 8 wherein the β1-integrin inhibitor is administered in 10. an amount effective to reduce the neurological deficits by at least about 80%.
- The method of claim 7 wherein the β 1-integrin inhibitor is a peptide 11. comprising a C-terminal LipAr motif.
- The method of claim 11 wherein the \beta1-integrin inhibitor is a peptide 12. comprising an amino acid sequence selected from the group consisting of WOPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAAIY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WOPPDADIY (SEO ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEQ ID NO:9), PRQAWRPIY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARIY (SEQ ID NO:12), RARIY (SEQ ID NO:13), ARIY (SEQ ID NO:14), and RIY.
- A method of treating a patient having a burn-type injury, the method 13. comprising administering a composition comprising a \$1-integrin inhibitor in an amount effective and over a period of time effective to reduce leukocyte-mediated tissue destruction.

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- The method of claim 13 wherein the β 1-integrin inhibitor is a peptide 14. comprising a C-terminal LipAr motif.
- The method of claim 14 wherein the β 1-integrin inhibitor is a peptide 15. comprising an amino acid sequence selected from the group consisting of WQPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAAIY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WQPPDADIY (SEQ ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEQ ID NO:9), PROAWRPIY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARIY (SEQ ID NO:12), RARIY (SEQ ID NO:13), ARIY (SEQ ID NO:14), and RIY.
- The method of claim 15 wherein the \beta1-integrin inhibitor is a peptide 16. comprising the amino acid sequence WQPPRARIY (SEQ ID NO:1).
- The method of claim 13 wherein the period of time is at least 1 hour. 17.
- The method of claim 17 wherein the period of time is at least 24 hours. 18.
- **19**. The method of claim 18 wherein the period of time is at least 48 hours.
- The method of claim 13 wherein the composition is administered 20. periodically over a predetermined period of time.
- 21. A method of treating a burn patient, the method comprising maintaining a composition comprising an effective amount of a \beta1-integrin inhibitor on a burn-type injury for a period of time effective to reduce leukocyte-mediated tissue destruction and achieve a desired degree of healing.



- The method of claim 21 wherein the β 1-integrin inhibitor is a peptide 22. comprising a C-terminal LipAr motif.
- The method of claim 22 wherein the β 1-integrin inhibitor is a peptide 23. comprising an amino acid sequence selected from the group consisting of WQPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAAIY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WQPPDADIY (SEQ ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEQ ID NO:9), PRQAWRPIY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARIY (SEQ ID NO:12), RARIY (SEQ ID NO:13), ARIY (SEQ ID NO:14), and RIY.
 - The method of claim 23 wherein the β 1-integrin inhibitor is a peptide 24. comprising the amino acid sequence WQPPRARIY (SEQ ID NO:1).
 - A method of treating a cancer patient, the method comprising administering 25. to the patient a composition comprising a \$1-integrin inhibitor in an amount effective to inhibit one or more of angiogenesis, cancer cell metastasis, cancer cell motility, or cancer cell migration.
 - The method of claim 25 wherein the β 1-integrin inhibitor is a peptide 26. comprising a C-terminal LipAr motif.
 - The method of claim 26 wherein the \beta1-integrin inhibitor is a peptide 27. comprising an amino acid sequence selected from the group consisting of WQPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAAIY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WQPPDADIY (SEQ ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEQ ID NO:9), PRQAWRPIY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARIY

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- 28. A method of treating a cancer patient, the method comprising administering to the patient a composition comprising a β1-integrin inhibitor in an amount effective to induce programmed cell death in cancerous tissue or restore normal cellular phenotype to cancerous tissue.
- 29. The method of claim 28 wherein the β1-integrin inhibitor is a peptide comprising a C-terminal LipAr motif.
- The method of claim 29 wherein the β1-integrin inhibitor is a peptide comprising an amino acid sequence selected from the group consisting of WQPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAAIY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WQPPDADIY (SEQ ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEQ ID NO:9), PRQAWRPIY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARIY (SEQ ID NO:12), RARIY (SEQ ID NO:13), ARIY (SEQ ID NO:14), and RIY.
- 31. The method of claim 28 further comprising administering a compound that inhibits the enzymatic degradation of the β1-integrin inhibitor.
- 32. A method of treating a patient for osteoporosis, the method comprising administering to the patient a composition comprising a β1-integrin inhibitor in an amount effective to inhibit osteoclast adhesion and bone resorption.
- 33. The method of claim 32 wherein the β1-integrin inhibitor is a peptide comprising a C-terminal LipAr motif.

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- A method of peripheralizing stem cells, the method comprising 35. administering to a patient a composition comprising a \$1-integrin inhibitor.
- The method of claim 35 wherein the β1-integrin inhibitor is a peptide **36.** comprising a C-terminal LipAr motif.
- The method of claim 36 wherein the \beta1-integrin inhibitor is a peptide 37. comprising an amino acid sequence selected from the group consisting of WOPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAATY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WQPPDADIY (SEQ ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEQ ID NO:9), PROAWRPTY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARTY (SEQ ID NO:12), RARIY (SEQ ID NO:13), ARIY (SEQ ID NO:14), and RIY.
- 38. A composition comprising \$1-integrin inhibitor and a pharmaceutically acceptable carrier.
- The composition of claim 38 wherein the \beta1-integrin inhibitor is a peptide 39. comprising a C-terminal LipAr motif.

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The composition of claim 39 wherein the β1-integrin inhibitor is a peptide comprising an amino acid sequence selected from the group consisting of WQPPRARIY (SEQ ID NO:1), WQPPRAAIY (SEQ ID NO:2), QPPRAAIY (SEQ ID NO:3), WQPPAARIY (SEQ ID NO:4), AQPPRARIY (SEQ ID NO:5), WAPPRARIY (SEQ ID NO:6), WQPPDADIY (SEQ ID NO:7), ARITGYIIY (SEQ ID NO:8), RARITGYIY (SEQ ID NO:9), PRQAWRPIY (SEQ ID NO:10), RPAPQRWIY (SEQ ID NO:11), PRARIY (SEQ ID NO:12), RARIY (SEQ ID NO:13), ARIY (SEQ ID NO:14), and RIY.